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ORIGINAL ARTICLE

Genome-wide association study across European and African American ancestries identifies a SNP in *DNMT3B* contributing to nicotine dependence

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Cigarette smoking is a leading cause of preventable mortality worldwide. Nicotine dependence, which reduces the likelihood of quitting smoking, is a heritable trait with firmly established associations with sequence variants in nicotine acetylcholine receptor genes and at other loci. To search for additional loci, we conducted a genome-wide association study (GWAS) meta-analysis of nicotine dependence, totaling 38,602 smokers (28,677 Europeans/European Americans and 9925 African Americans) across 15 studies. In this largest-ever GWAS meta-analysis for nicotine dependence and the largest-ever cross-ancestry GWAS meta-analysis for any smoking phenotype, we reconfirmed the well-known *CHRNA5-CHRNA3-CHRNA4* genes and further yielded a novel association in the DNA methyltransferase gene *DNMT3B*. The intronic *DNMT3B* rs910083-C allele (frequency = 44–77%) was associated with increased risk of nicotine dependence at $P = 3.7 \times 10^{-8}$ (odds ratio (OR) = 1.06 and 95% confidence interval (CI) = 1.04–1.07 for severe vs mild dependence). The association was independently confirmed in the UK Biobank ($N = 48,931$) using heavy vs never smoking as a proxy phenotype ($P = 3.6 \times 10^{-4}$, OR = 1.05, and 95% CI = 1.02–1.08). Rs910083-C is also associated with increased risk of squamous cell lung carcinoma in the International Lung Cancer Consortium ($N = 60,586$, meta-analysis $P = 0.0095$, OR = 1.05, and 95% CI = 1.01–1.09). Moreover, rs910083-C was implicated as a *cis*-methylation quantitative trait locus (QTL) variant associated with higher *DNMT3B* methylation in fetal brain ($N = 166$, $P = 2.3 \times 10^{-26}$) and a *cis*-expression QTL variant associated with higher *DNMT3B* expression in adult cerebellum from the Genotype-Tissue Expression project ($N = 103$, $P = 3.0 \times 10^{-6}$) and the independent Brain eQTL Almanac ($N = 134$, $P = 0.028$). This novel *DNMT3B* *cis*-acting QTL variant highlights the importance of genetically influenced regulation in brain on the risks of nicotine dependence, heavy smoking and consequent lung cancer.

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INTRODUCTION

Cigarette smoking is a leading cause of preventable death, resulting annually in nearly 6 million premature deaths worldwide.¹ Smoking-related deaths are most often attributed to

increased rates of cancer, cardiovascular disease and chronic obstructive pulmonary disease.² Despite the well-known adverse health effects, an estimated 45.3 million US adults smoke cigarettes, of whom over 68% report wanting to quit.^{3,4}

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Cigarette smoking is a complex multi-step behavior involving initiation, regular smoking, nicotine dependence, cessation and relapse. Some regular smokers maintain low-level smoking without developing symptoms of dependence,⁵ while others become heavily dependent smokers and experience the most difficulty with cessation and the highest risk of relapse.^{6,7} Nicotine dependence has high heritability (estimates up to 75%),^{8,9} and besides reducing the likelihood of quitting smoking, it is predictive of withdrawal severity,¹⁰ response to treatment¹¹ and smoking-related health outcomes.^{12,13}

Genome-wide association study (GWAS) analyses of nicotine dependence phenotypes^{14–24} have firmly established associations with several loci, including nicotinic acetylcholine receptor genes on chromosomes 15q25 (*CHRNA5-CHRNA3-CHRNA4*), 8p11 (*CHRNA3-CHRNA6*) and 20q13 (*CHRNA4*). The largest GWAS meta-analyses relied on widely ascertained phenotypes such as cigarettes per day (CPD),^{16–18} which represents only one of several components of nicotine dependence.²⁵ Focusing GWAS on nicotine dependence rather than CPD may improve statistical power for identifying variants that influence the broader construct of dependence.¹⁹ This idea is supported by our prior nicotine dependence GWAS meta-analysis (total $N=17,074$ ever-smokers of European/European American ancestry (EUR)) that discovered associations with *CHRNA4* single-nucleotide polymorphisms (SNPs) that were driven by time to first cigarette in the morning (TTFC) and had not been detected in GWAS meta-analyses of CPD with much larger sample sizes.²³ To improve statistical power further and to search for additional loci, we more than doubled our sample size to perform the largest GWAS meta-analysis of nicotine dependence to date, including 38,602 ever-smokers (28,677 of EUR and 9925 of African American (AA) ancestries) across 15 studies. We extended our study to include correlations with DNA methylation (DNAm) and RNA expression (RNAexp) of nearby genes across human brain tissues and evaluated associations with a critical smoking-related outcome: lung cancer (total $N=81,821$ cases and controls).

MATERIALS AND METHODS

Study protocols received institutional review board approval at their respective sites. All study participants provided written informed consent.

We included the five studies from our prior GWAS meta-analysis²³ and 10 additional studies. Details of their study design, genotyping, quality control, 1000 Genomes (1000G) imputation and analysis are provided in Supplementary Methods and Supplementary Table 1.

Nicotine dependence phenotype

We included studies with SNP genotypes and Fagerström Test for Nicotine Dependence (FTND) data²⁶ collected among smokers. FTND scores range from 0 (no dependence) to 10 (highest dependence level). As before,²³ we used FTND to categorize nicotine dependence as mild (scores 0–3), moderate (scores 4–6) or severe (scores 7–10). Two of the 15 studies additionally included low-intensity smokers who reported CPD as ≤ 10 but had no data available on the other FTND items and were defined as mildly dependent. Concordance rates between these FTND and CPD categories showed minimal phenotype misclassification (Supplementary Methods).

Nicotine dependence GWAS meta-analysis and independent follow-up

We used linear regression to test SNP/indel associations with categorical nicotine dependence (mild = 0/moderate = 1/severe = 2) in each separate study and ancestry group. Covariates included age, sex, principal components and study-specific covariates (as needed); additional adjustment for family structure was made in studies with relatives included (Supplementary Methods).

We combined GWAS results, using METAL²⁷ with fixed-effects inverse variance-weighting meta-analysis, across all studies with FTND data to maximize statistical power. Genomic control was applied to the deCODE results to adjust for inflation due to relatedness among participants; all

other studies had low inflation values ($\lambda \leq 1.02$). We excluded SNPs/indels with minor allele frequency $< 1\%$ in 1000G EUR or African (AFR) panels, depending on the ancestry group analyzed. The standard threshold ($P < 5 \times 10^{-8}$), originally based on 1 million independent tests genome-wide as computed using HapMap-based imputation for EUR studies,²⁸ has been validated for 1000G-imputed GWAS of common variants.²⁹ Rather than imposing a more stringent, yet to be consistently determined, threshold when analyzing common variants across EUR and AA studies,^{29–32} we carried forward novel variation implicated at $P < 5 \times 10^{-8}$ and relied on confirmation in an independent study to declare genome-wide significance. For this confirmation step, we utilized UK Biobank ($N=48,931$ EUR participants) results with heavy, defined as pack-years ($[\text{CPD}/20 \text{ cigarettes per pack}] \times \text{years smoked}$) ≥ 10 , vs never smoking as a proxy phenotype.²⁴ This prior GWAS was designed as a nested case-control analysis that sampled the extremes of smoking behavior, and thus did not encompass light smoking.

Regional association plots were created using LocusZoom³³ with linkage disequilibrium (LD) estimates of r^2 and D' based on 1000G EUR and AFR panels. Allele frequencies were weighted by sample size. Odds ratio (OR) estimates were computed using the β estimate from the SNP term in the linear regression model ($e^{[2 \times \beta]}$ for severe vs mild dependence) and then compared across studies using the Forest Plot Viewer.³⁴ Heterogeneity across studies was assessed using the I^2 index.³⁵

SNP associations with DNAm and RNAexp

The top novel nicotine dependence-associated variant was assessed for *cis*-acting regulatory effects, using previously published methylation quantitative trait locus (meQTL) and expression QTL (eQTL) studies of postmortem brain tissues (Supplementary Methods). We used meQTL results from 166 fetal brain samples in the Human Developmental Biology Resource and UK Medical Research Council Brain Banks network; these samples were not dissected further into different tissues.³⁶ SNP genotypes were imputed using 1000G, and DNAm was measured using the Illumina HumanMethylation450 BeadChip. The QTL results were obtained at <http://epigenetics.essex.ac.uk/meQTL/>.

We assessed SNP associations with RNAexp of nearby genes (± 1 Mb around the transcription start sites) across adult brain tissues from the Genotype-Tissue Expression (GTEx; <https://gtexportal.org>) project³⁷ with 1000G-imputed genotypes and RNA sequencing measures from up to 103 deceased persons. We followed up *cis*-eQTL SNP results in the Brain eQTL Almanac data set ($N=134$ EUR deceased persons, <http://www.braineac.org/>), using RNAexp probes measured on Affymetrix Human Exon 1.0 ST arrays across brain tissues, many of which overlap with tissues available in GTEx.³⁸

SNP associations with lung cancer

The top novel nicotine dependence-associated variant was tested for association with lung cancer using 27,349 cases and 54,472 controls from > 30 EUR studies in the Transdisciplinary Research for Cancer in Lung of the International Lung Cancer Consortium (TRICL-ILCCO). Their genotyping, quality control, 1000G imputation and statistical analyses were previously described.³⁹ Briefly, the SNP associations were derived from a meta-analysis of logistic regression model results that compared lung cancer cases and controls, adjusting for age, sex and the first two principal components. Histological subtypes of adenocarcinoma and squamous cell lung carcinoma cases were compared, separately, to controls. We removed the EAGLE study from the original set of lung cancer studies to avoid overlap in the nicotine dependence and lung cancer meta-analyses. All lung cancer analyses included ever- and never-smokers; our follow-up association testing made additional adjustments for smoking history (ever vs never) and pack-years.

RESULTS

We performed GWAS analyses across 15 studies, totaling 38,602 (28,677 EUR and 9925 AA) ever-smokers (Supplementary Table 2), with nicotine dependence defined as mild ($N=17,796$; 46.1%), moderate ($N=13,527$; 35.0%) or severe ($N=7279$; 18.9%). More than 99% of the participants were ≥ 18 years old. Males constituted 53.2% of the total sample size.

Our GWAS meta-analysis tested nearly 18 million genotyped and 1000G-imputed SNPs/indels for association with mild/

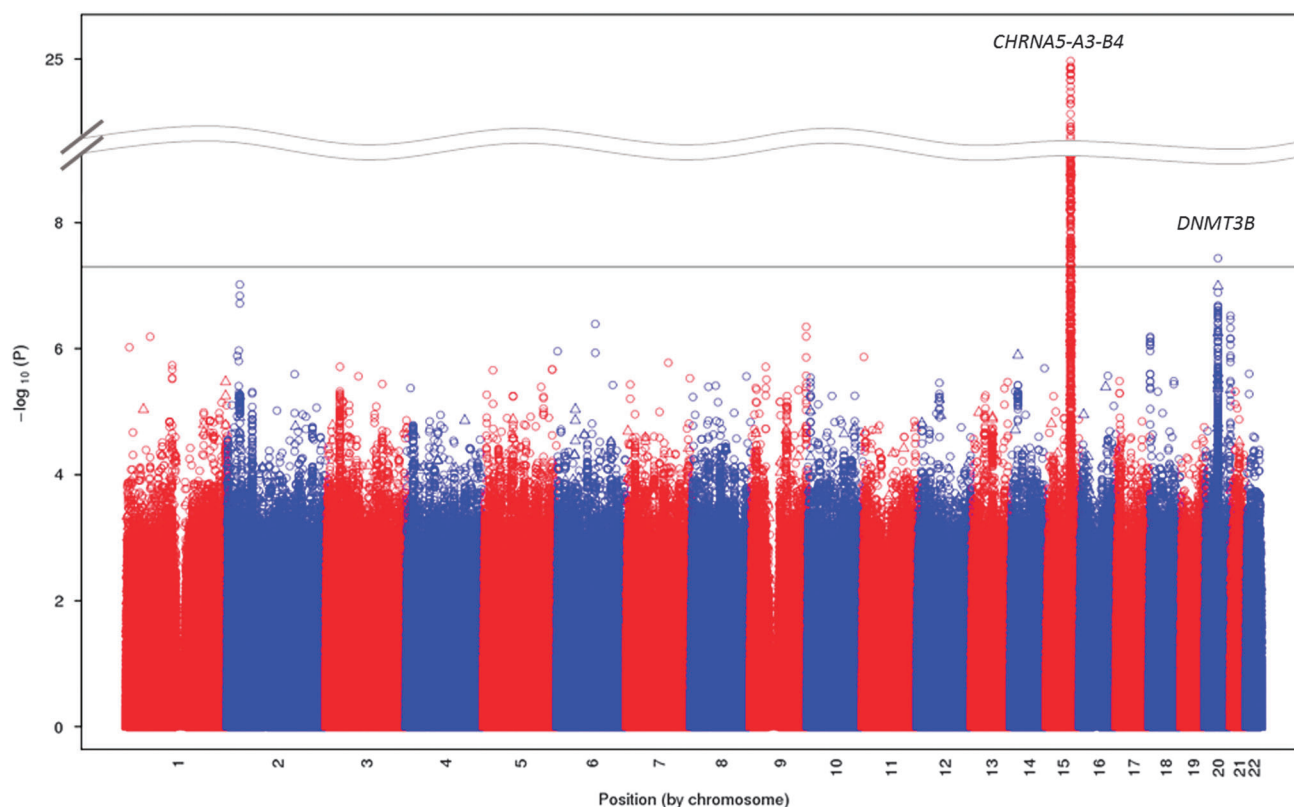


Figure 1. Manhattan plot of SNP and indel associations with nicotine dependence from GWAS meta-analysis across 15 studies (total $N = 38,602$ European/European Americans and African Americans). The $-\log_{10}$ meta-analysis P -values are plotted by chromosomal position of SNPs (depicted as circles) and indels (depicted as triangles). The genome-wide statistical significance threshold ($P < 5 \times 10^{-8}$) is shown as a solid black line. GWAS, genome-wide association study; SNP, single-nucleotide polymorphism.

moderate/severe dependence ($\lambda = 1.027$, Supplementary Figure 1A). We observed SNP/indel associations at meta-analysis $P < 5 \times 10^{-8}$ at two loci (Figure 1): the well-known chromosome 15q25 locus spanning *CHRNA5-CHRNA3-CHRNA4* (smallest meta-analysis $P = 1.3 \times 10^{-25}$) and a novel chromosome 20q11 locus spanning the DNA (cytosine-5)-methyltransferase 3 beta (*DNMT3B*) gene (smallest meta-analysis $P = 3.7 \times 10^{-8}$). No additional loci were identified at $P < 5 \times 10^{-8}$ in the ancestry-specific GWAS meta-analyses (Supplementary Figures 1B–C and 2A–B).

DNMT3B SNP associations with nicotine dependence

Rs910083, an intronic *DNMT3B* SNP, was identified across both ancestries: meta-analysis $P = 3.7 \times 10^{-8}$ and β (standard error) = 0.032 (0.0057) for the C allele (Table 1), corresponding to OR (95% confidence interval (CI)) of 1.06 (1.04–1.07) for severe vs mild dependence (Supplementary Figure 3). Rs910083 was imputed well (quality scores = 0.98–1), and it showed no evidence of heterogeneity across studies ($P = 0.71$).

Rs910083-C, the minor allele for EUR (frequency = 44%) but the major allele for AA (frequency = 77%), is associated with increased nicotine dependence risk (Table 1): EUR-specific meta-analysis $P = 4.1 \times 10^{-5}$ and OR (95% CI) = 1.06 (1.03–1.08); and AA-specific meta-analysis $P = 7.3 \times 10^{-5}$ and OR (95% CI) = 1.10 (1.05–1.15). Many SNPs/indels were in moderate to high LD ($r^2 > 0.4$) with rs910083 in 1000G EUR, spanning 220 kb (chr20:31,268,924–31,488,466) and including *DNMT3B* and its neighboring genes, microtubule-associated protein, RP/EB family, member 1 (*MAPRE1*) and COMM domain containing 7 (*COMM7*). However, in 1000G AFR, SNPs in LD ($r^2 > 0.4$) with rs910083 were localized to a 47 kb

region (chr20:31,356,560–31,403,394) including only *DNMT3B* (Figures 2A–B, NCBI build 37 positions).

No *DNMT3B* variants have been implicated previously for any substance use disorder (SUD) phenotype. Upstream of *DNMT3B*, chromosome 20q11 also harbors the nucleolar protein 4-like (*NOL4L*) gene, which was reported at genome-wide significance for heavy vs never smoking in the UK Biobank for the indel rs57342388.²⁴ This indel was associated at meta-analysis $P = 0.0017$ in our study (Table 1): OR (95% CI) = 1.04 (1.02–1.07) for severe vs mild dependence for the insertion allele, consistent with the prior result. Rs57342388 is located 216 kb upstream of our top *DNMT3B* SNP rs910083. The two variants are weakly correlated ($r^2 = 0.11$ in 1000G EUR where minor allele frequency = 2% for rs57342388 vs 18% for rs910083, $r^2 = 0.0022$ in AFR where minor allele frequency = 19% vs 42%) but are in moderate to high LD ($D' = 0.57$ in EUR, $D' = 1$ in AFR). In follow-up testing with both SNPs included in the same model, both were associated with nicotine dependence (meta-analysis $P = 1.7 \times 10^{-6}$ for rs910083 and 8.3×10^{-3} for rs57342388), showing that our observed *DNMT3B* association signal is not explained by the previously reported *NOL4L* signal.

We tested rs910083-C for association with each of the specific FTND items, as presented in Supplementary Table 3. The rs910083-C association was driven most strongly by TTFC (meta-analysis $P = 1.2 \times 10^{-4}$). Its next most significantly associated FTND item was CPD (meta-analysis $P = 0.0011$). TTFC is an indicator of withdrawal severity upon awakening⁴⁰ and behavioral automaticity (habitual smoking without awareness or cognitive control⁷). Although TTFC has its distinct features (for example, strongest

Table 1. SNP associations in the novel *DNMT3B* gene and previously reported genes that were identified at $P < 5 \times 10^{-7}$ in the nicotine dependence GWAS meta-analysis

Chr	Gene/nearest gene	Prior GWAS reported smoking phenotype	SNP (allele)	Nicotine dependence meta-analyses among ever regular smokers						UK Biobank heavy vs never smokers (N = 48,931) ²⁴	
				EUR studies (N = 28,677)			AA studies (N = 9925)			EUR and AA studies (N = 38,602)	
				β (SE)	P	β (SE)	β (SE)	P	β (SE)	β (SE)	P ^a
20q11	<i>DNMT3B</i>	None	rs910083 (C)	0.027 (0.0065)	4.2×10^{-5}	0.047 (0.012)	0.032 (0.0057)	7.3×10^{-5}	0.048 (0.014)	3.6×10^{-4}	
	<i>NOL4L</i>	Heavy vs never smoking ²⁴	rs57342388 (CACGG)	0.021 (0.0075)	0.0059	0.032 (0.020)	0.022 (0.0070)	0.10	0.094 (0.016)	4.7×10^{-9}	
	<i>CHRNA4</i>	None	rs6062901 ^b (G)	0.043 (0.0088)	8.3×10^{-7}	0.022 (0.010)	0.034 (0.0067)	0.035	0.048 (0.018)	0.0075	
20q13		Nicotine dependence ²³	rs2273500 ^b (C)	0.046 (0.0098)	2.0×10^{-6}	0.0085 (0.016)	0.036 (0.0083)	0.59	0.091 (0.019)	1.4×10^{-6}	
		Heavy vs never smoking ²⁴	rs11697662 ^b (C)	0.033 (0.0085)	9.4×10^{-5}	0.013 (0.011)	0.025 (0.0066)	0.20	0.091 (0.017)	1.0×10^{-7}	
		None	rs56116178 ^c (G)	0.053 (0.010)	4.5×10^{-7}	NA	0.053 (0.010)	NA	0.097 (0.022)	9.3×10^{-6}	
9q34	<i>DBH</i>	Heavy vs never smoking ²⁴	rs111280114 ^c (G)	0.052 (0.010)	6.4×10^{-7}	NA	0.052 (0.0098)	NA	0.099 (0.022)	6.0×10^{-6}	
		Smoking cessation ¹⁷	rs3025343 ^c (A)	0.043 (0.0099)	1.7×10^{-5}	NA	0.042 (0.0098)	NA	0.091 (0.021)	1.1×10^{-5}	

Abbreviations: AA, African American; EUR, European/American; GWAS, genome-wide association study; LD, linkage disequilibrium; SE, standard error; SNP, single-nucleotide polymorphism. The SNP with the smallest nicotine dependence GWAS meta-analysis P is shown, along with nearby SNPs previously reported in other GWAS of smoking phenotypes. Results from the UK Biobank heavy vs never smoking GWAS are also presented. NA, not available due to minor allele frequency < 1%. ^a P -values correspond to genomic control being applied. ^bRs2773500 and rs11697662 were originally implicated in EUR studies, where they exist in moderate LD ($r^2 = 0.64$ and $D' = 0.95$ in 1000G EUR). Rs6062901 had $r^2 = 0.47$ – 0.57 and $D' = 0.73$ – 0.84 with rs2273500 and rs11697662 in 1000G EUR and $r^2 = 0.72$ – 0.92 in 1000G AFR. ^crs56116178 and rs111280114 had $r^2 = 0.98$ and $D' = 1$ with one another and $r^2 = 0.72$ – 0.73 and $D' = 0.96$ with rs3025343 in 1000G EUR.

predictor of cessation success among the FTND items), it is correlated with heaviness of smoking as captured by CPD.⁷

DNMT3B SNP association with heavy smoking in an independent study

Using heavy vs never smoking GWAS results from the UK Biobank ($N = 24,457$ heavy and 24,474 never smokers),²⁴ we found that rs910083-C is associated at $P = 3.6 \times 10^{-4}$ and OR (95% CI) = 1.05 (1.02–1.08) for heavier smoking. Although effect sizes were small, its associations with risks of nicotine dependence and heavy smoking were consistent (Table 1).

DNMT3B SNP associations with DNAm and RNAexp

Using a genome-wide meQTL study of 166 fetal brain samples,³⁶ we found that rs910083-C associates with higher DNAm levels of the probe cg13636640, located 252 base pairs upstream of the *DNMT3B* gene ($\beta = 0.082$, $P = 2.3 \times 10^{-26}$). This probe was the only one observed in the region with significant mQTL variants (Supplementary Figure 4). There were a total of 140 SNPs associated with DNAm of this probe (smallest $P = 1.8 \times 10^{-45}$), and we observed associations of these SNPs with nicotine dependence with meta-analysis P -values ranging from 3.66×10^{-8} for rs910083 to 0.0051 (Supplementary Table 4). To our knowledge, there are no similar data to assess *cis*-meQTL effects in brain tissues from adults with no psychiatric disease. However, we assessed the *cis*-acting effect of rs910083 on RNAexp across several adult brain tissues using two independent data sets, GTEx³⁷ followed by Brain eQTL Almanac.³⁸

Across the 13 brain tissues in GTEx, we observed the highest *DNMT3B* gene expression levels in cerebellum (median log₁₀ RPKM = 0.304) and cerebellar hemisphere (median log₁₀ RPKM = 0.298), compared to median log₁₀ RPKM ≤ -0.325 for all other brain tissues (Supplementary Figure 5). Moreover, across the brain tissues, rs910083 was most significantly associated with *DNMT3B* gene expression levels in cerebellum ($P = 3.0 \times 10^{-6}$) and cerebellar hemisphere ($P = 7.0 \times 10^{-7}$), with the C allele associated with higher *DNMT3B* expression (Figure 3 and Supplementary Table 5). We replicated this pattern in the Brain eQTL Almanac, where *DNMT3B* mRNA transcript expression levels were highest in cerebellar cortex (Supplementary Figure 6)—the outer layer of the cerebellum that comprises most of its volume. Consistent with GTEx, rs910083-C was associated with increased *DNMT3B* mRNA transcript expression levels specifically in cerebellar cortex ($P = 0.028$).

Beyond the brain tissues in GTEx, rs910083 is associated with RNAexp of other genes within 1 MB (Supplementary Table 6): *MAPRE1* (smallest $P = 7.3 \times 10^{-17}$ in sun exposed skin), *COMMD7* (smallest $P = 1.0 \times 10^{-6}$ in colon) and BPI fold containing family B, member 2 (*BPIFB4*, smallest $P = 5.4 \times 10^{-5}$ in artery).

DNMT3B SNP associations with lung cancer

We assessed rs910083 for association with lung cancer using a GWAS meta-analysis of EUR studies from TRICL-ILCCO³⁹ ($N = 27,349$ cases and 54,472 controls, Supplementary Table 7). Rs910083-C is significantly associated with increased risk of squamous cell carcinoma ($N = 6937$ cases and 53,649 controls, meta-analysis $P = 0.0095$ and OR [95% CI] = 1.05 [1.01–1.09]), consistent with the increased nicotine dependence risk. Rs910083 is not associated with adenocarcinoma.

We evaluated the effect of adjusting for smoking on the rs910083 association with squamous cell carcinoma in the studies with smoking data readily available. We found that the level of statistical significance and magnitude of association were both weakened with adjustment for ever/never smoking and pack-years (Supplementary Table 8), suggesting that the rs910083

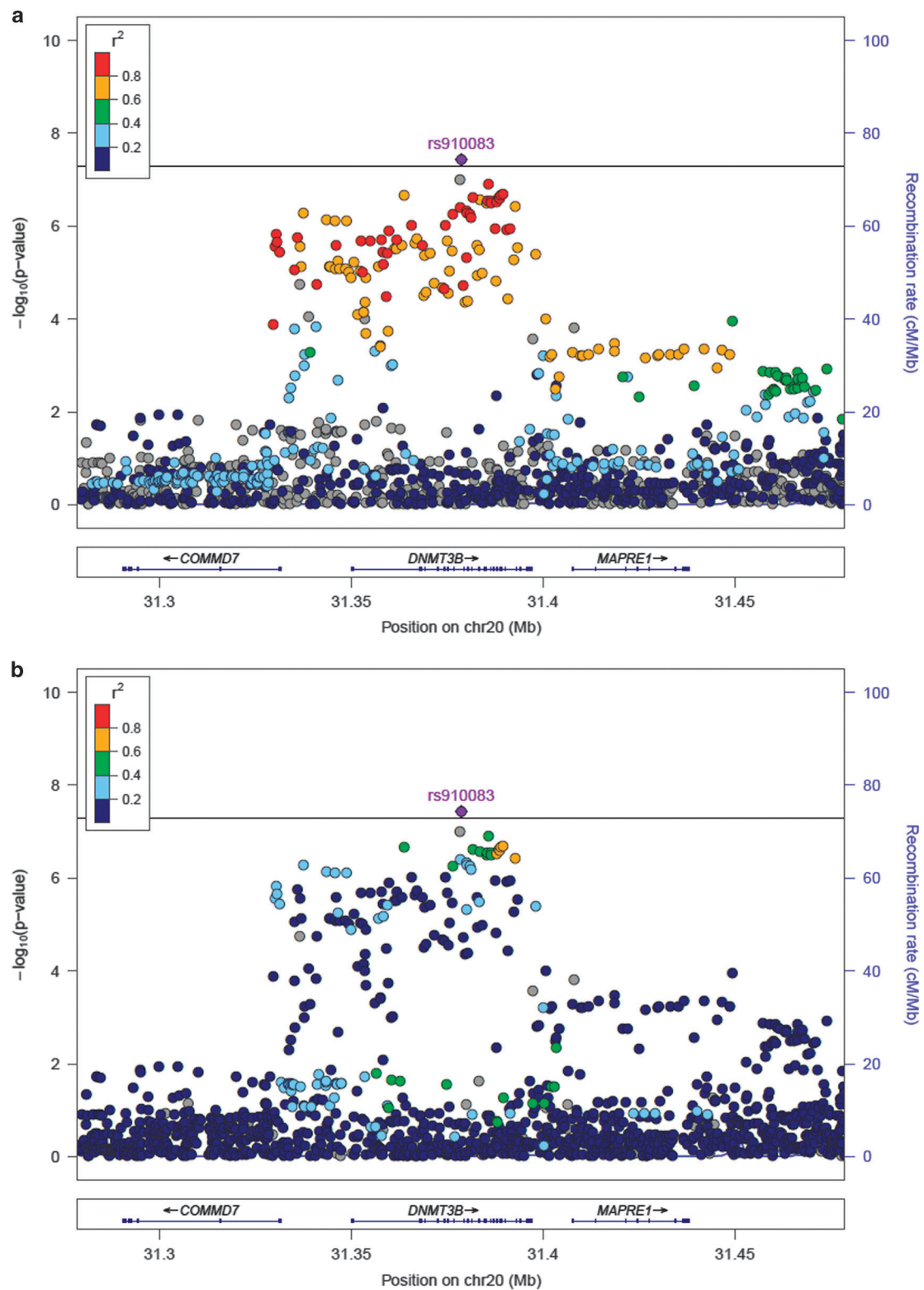


Figure 2. Novel *DNMT3B* SNP associations with nicotine dependence from GWAS meta-analysis of EUR and AA studies. SNP and indel associations are shown across *DNMT3B* and its 100 kb flanking region (NCBI build 37 positions presented). r^2 values between the top SNP rs910083 and all other SNPs are shown in reference to 1000 Genomes panels: (a) EUR and (b) AFR. Indels with missing r^2 values are indicated in gray. The P -value threshold of 5×10^{-8} is marked by the solid black line. AA, African American; EUR, European/European American; GWAS, genome-wide association study; SNP, single-nucleotide polymorphism.

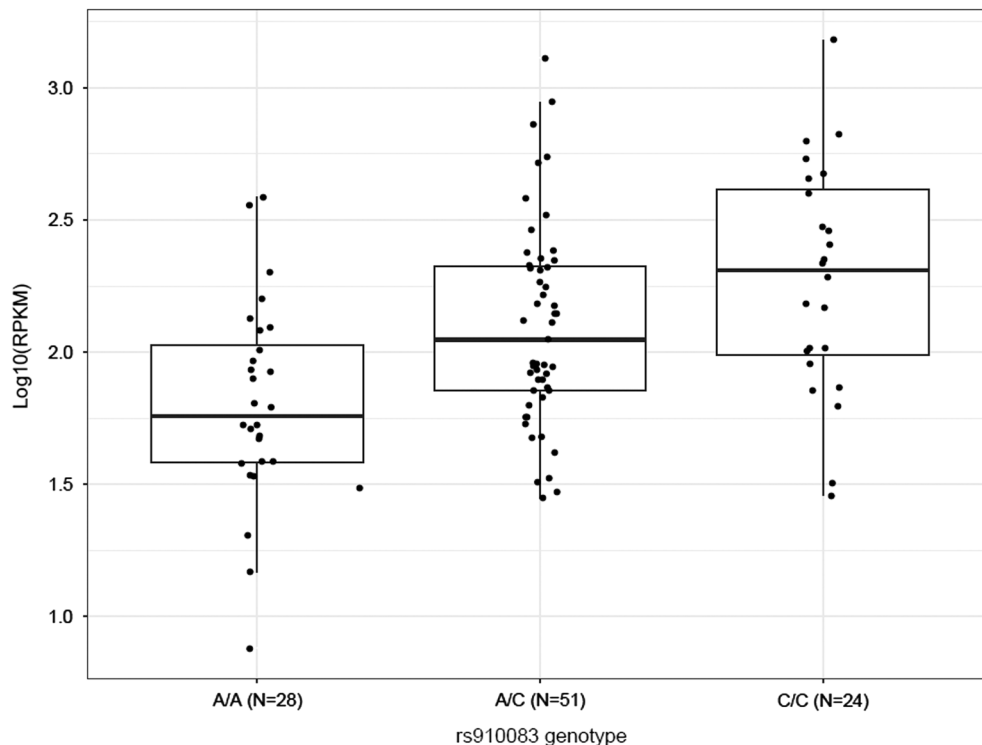


Figure 3. Normalized *DNMT3B* gene expression levels as a function of rs910083 genotype in cerebellum from the Genotype-Tissue Expression (GTEx) project. The box lines mark the first quartile, median and third quartile; and the whiskers are marked by the highest and lowest data points within the 1.5 × inter-quartile range (third–first quartile) to show outliers that fall outside of these boundaries.

association with squamous cell carcinoma is mediated by smoking.

DISCUSSION

This largest-ever GWAS meta-analysis for nicotine dependence, and the largest-ever cross-ancestry GWAS meta-analysis for any smoking phenotype, identified rs910083 as a novel SNP that regulates *DNMT3B* in human brain and contributes to risks of nicotine dependence and heavy smoking. Rs910083 was discovered via meta-analysis of two ancestry groups; the association signal includes SNPs in high LD with rs910083 across the *COMMD7*, *DNMT3B* and *MAPRE1* genes in EUR ancestry, but LD is localized to the *DNMT3B* gene in AA ancestry. Moreover, rs910083 was implicated as a *cis*-acting QTL SNP that influences *DNMT3B* DNAm in fetal brain and *DNMT3B* RNAexp in adult cerebellum, with the C allele being associated with higher DNAm and RNAexp levels. While this pattern might contrast the traditional view of higher DNAm being correlated with lower RNAexp, the observed effects reflect temporal and spatial differences, thus limiting our ability to draw direct correlations. Nonetheless, genome-wide QTL comparisons in human brain have shown that almost half of SNPs that act as both an eQTL and meQTL show the same direction of association for DNAm and RNAexp,⁴¹ as we observed here for *DNMT3B* and before for *CHRNA5*.⁴² The previously established nicotinic acetylcholine receptor genes also harbor nicotine dependence-associated SNPs with important consequences for gene regulation, including noncoding SNPs that correlate with DNAm,⁴² splicing,²³ and/or RNAexp^{43,44} in brain tissues that are frequently studied for nicotine and other SUDs because of their role in primary reward pathways and executive function, such as prefrontal cortex.⁴⁵ This newly identified *DNMT3B* SNP association highlights changes in DNAm in fetal brain and with RNAexp specifically in cerebellum, a part of the brain that has often been

overlooked despite some indications for its involvement in the neurobiology of addiction.^{46–48}

DNMT3B encodes a DNA methyltransferase prominently involved in *de novo* DNAm that establishes patterns early in development; it may also contribute to maintenance DNAm.⁴⁹ Although initially reported for maintaining DNAm at imprinted loci,⁵⁰ later evidence showed that other members of the DNMT family (*DNMT3A* and *DNMT3L*) are required for imprinting, while *DNMT3B* may not play an essential role.⁵¹ Mouse models have shown that complete loss of *DNMT3B* function is embryonically lethal.⁵⁰ However, recessive inheritance of rare mutations that render *DNMT3B* partially functional are known to cause Immunodeficiency, Centromeric instability and Facial dysmorphism syndrome, which manifests with growth and neurodevelopmental abnormalities.⁴⁹ Because of the critical role that *DNMT3B* plays in establishing methylation, altered expression has been associated with Immunodeficiency, Centromeric instability and Facial dysmorphism syndrome at >700 genes involved in brain development and other processes.⁵² These and other genes that are regulated by *DNMT3B* methylation represent candidate genes that may directly contribute to nicotine dependence susceptibility.

DNMT3B has not previously been connected with the biology underlying the risk of nicotine dependence or any other SUD. However, there is evidence that *in vitro* cigarette smoke exposure leads to increased *DNMT3B* expression in human respiratory epithelial cells, and *DNMT3B* overexpression results in downstream hypermethylation that has been widely implicated in lung cancer.⁵³ Given this indication, DNMT inhibition has been an active area of research for cancer treatment; two inhibitor agents are currently approved by the US Food and Drug Administration (decitabine, which shows high affinity for *DNMT3A/3B* over *DNMT1*,⁵⁴ and azacytidine), and at least one other promising agent (zebularine) awaits clinical trial testing.⁵⁵ With the *DNMT3B* variant discovery for nicotine dependence, DNMT3B inhibition

may merit future study for smoking cessation treatment. Rs910083 resides in an active promoter marked by H3K9ac in several brain regions examined in the Roadmap Epigenomics Project,⁵⁶ and it associates with *DNMT3B* RNAexp specifically in the cerebellum. This finding does not negate the importance of other brain regions known to be involved in SUDs but brings up the possibility of altered gene regulation in the cerebellum contributing to the complex neurobiological pathway leading to dependence. A main function of the cerebellum is motor coordination, but it is also involved in non-motor functions relevant to SUD, including reward.^{46–48} The cerebellum responds to acute and long-term exposures to nicotine^{57–62} and other substances,⁴⁷ and it makes functional connections with the prefrontal cortex and other brain tissues that are widely recognized for their involvement in SUDs.⁴⁷

Our discovery of the rs910083 association with nicotine dependence was made possible by assembling the largest possible sample size of FTND studies comprised of ever-smokers to maximize statistical power. No large FTND studies were left for replication, but because CPD is a central component of both the FTND and the heavy vs never smoking definition used in the UK Biobank, we tested for independent confirmation of the novel association with heavy vs never smoking and found that rs910083-C also conferred risk in the UK Biobank.

Larger GWAS meta-analyses have been reported for other smoking phenotypes such as ever vs never smoking, but these studies were comprised only of EUR participants and based on HapMap imputation.^{16–18} Rs910083, a 1000G-imputed SNP, was not captured in these studies, but 9 of the 18 *DNMT3B* SNPs associated with nicotine dependence at meta-analysis $P < 5 \times 10^{-7}$ in the present study were HapMap-imputed (Supplementary Table 9); these SNPs were in strong LD with rs910083 among EURs ($r^2 = 0.78–0.99$ in 1000G EUR) but weaker LD among AAs ($r^2 = 0.29–0.76$ in 1000G AFR). Using results from the largest GWAS meta-analysis of CPD (Tobacco and Genetics (TAG) consortium, $N = 38,181$ EUR ever smokers independent of the ones included here), we found that the nine HapMap-imputed *DNMT3B* SNPs were associated with CPD at P -values ranging from 0.027 to 0.059 and a consistent direction of association with nicotine dependence; in comparison, P -values for ever vs never smoking ($N = 74,035$ in the TAG consortium) ranged from 0.049 to 0.34 (Supplementary Table 9). We caution that the best *DNMT3B* signal in the TAG consortium was observed for CPD at only nominal significance (smallest $P = 0.027$), despite having a nearly equivalent sample size as our study. However, our study yielding more statistically significant *DNMT3B* SNP associations with nicotine dependence (smallest $P = 3.7 \times 10^{-8}$) is likely due to a combination of factors, including (1) reliance on FTND, a multi-dimensional phenotype that encompasses CPD and other important features of smoking behavior including TTFC that drove the rs910083 association, and (2) 1000G imputation which has been shown to strengthen association signals for some loci due to the finer mapping available.^{63,64} We have similarly observed more statistically significant associations with nicotine dependence and stronger effect sizes, compared to CPD, in prior studies of *CHRNA3*¹⁹ and *CHRNA4*.²³

Until now, the only common *DNMT3B* variant implicated by GWAS was identified in a study of inflammatory bowel disease (intronic rs4911259).⁶⁵ This SNP was associated with nicotine dependence in our meta-analysis ($P = 3.5 \times 10^{-6}$) and is in LD with our top SNP rs910083 ($r^2 = 0.76$ and $D' = 1$ in 1000G EUR, $r^2 = 0.16$ and $D' = 1$ in 1000G AFR). Cigarette smoking is the environmental factor most consistently associated with inflammatory bowel disease,⁶⁶ and these shared SNP association signals suggest that *DNMT3B* SNPs may exert pleiotropic effects. Alternatively, it is possible that smoking mediates the *DNMT3B* SNP association with inflammatory bowel disease, but the inflammatory bowel disease GWAS did not include adjustment for smoking.⁶⁵

Beyond finding *DNMT3B* and (as expected) *CHRNA5-CHRNA3-CHRNA4*, our GWAS meta-analysis resulted in $P < 5 \times 10^{-7}$ for two other loci previously implicated in smoking, *CHRNA4* on chromosome 20q13^{23,24} and dopamine β -hydroxylase (*DBH*) on chromosome 9q34.^{17,24} Our prior nicotine dependence GWAS meta-analysis of EUR studies identified the *CHRNA4* splice site SNP rs2773500.²³ Rare *CHRNA4* variants have also been found to associate with nicotine dependence.^{67,68} Our study supported common *CHRNA4* SNP associations among EUR samples, but no association was detected for these SNPs among AA studies (Table 1).

DBH is a strong functional candidate for influencing nicotine dependence. The dopaminergic system lies at the core of the brain's reward pathway, and the *DBH* enzyme converts dopamine into norepinephrine. An upstream *DBH* SNP (rs3025343) was identified in a GWAS of smoking cessation (current vs former smokers)^{16–18} and later independently replicated.^{69,70} Consistent with rs3025343-A being associated with reduced success of quitting smoking, its phenotypic profile has been expanded to include associations with: (1) heavier smoking ($N = 48,931$ in the UK Biobank, $P = 1.2 \times 10^{-5}$)²⁴; (2) higher FTND scores ($N = 1430$ EUR participants, $P = 0.023$)⁷¹; and (3) higher nicotine dependence risk in our EUR studies ($N = 28,677$, meta-analysis $P = 1.7 \times 10^{-5}$). Smaller P -values were found for other 1000G-imputed upstream *DBH* SNPs in the UK Biobank and our study (rs111280114 and rs56116178, respectively; Table 1); the minor alleles of these SNPs were similarly associated with increased risks among EUR studies (Supplementary Figure 7 for rs56116178). Because these *DBH* SNPs all occur at $< 1\%$ frequency among AAs, studying *DBH* variation on nicotine dependence risk in this ancestry group will require larger sample sizes or an alternative study design.

Nicotine dependence-associated variants in *CHRNA5-CHRNA3-CHRNA4*^{44,72} and *CHRNA4*^{23,67} have been previously shown to associate with lung cancer and other smoking-related diseases. Our study shows that the nicotine dependence-associated SNPs in *DNMT3B* and *DBH* are also associated with lung cancer (Supplementary Table 7). These findings may reflect the SNPs acting indirectly on lung through their influence on smoking (Supplementary Table 8). Alternatively, because *DBH* is expressed in the lung⁷³ and *DNMT3B* overexpression has been shown in lung cancer, we cannot exclude the possibility that either of these SNPs act directly to promote lung cancer through an unknown mechanism.⁵³ The *DNMT3B* and *DBH* SNPs were both associated with squamous cell lung carcinoma. This histological subtype has a strong association with smoking and occurs infrequently in never-smokers. In contrast, neither SNP is associated with adenocarcinoma, a subtype that has a weaker association with smoking⁷⁴ and an increasing prevalence over time among never-smokers.⁷⁵ Histology-specific associations are not uncommon for lung cancer genetic loci.³⁹

Our findings expand the known genetic architecture of nicotine dependence, by showing that the *DNMT3B* SNP rs910083 increases the likelihood of developing nicotine dependence as observed across two different ancestries, smoking heavily, and consequently incurring a heightened risk of lung cancer.^{53,55} The convergence of prior and current findings indicate that the complex neurobiology underlying nicotine dependence involves several sequence variants with functional and regulatory effects across distinct brain tissues.

CONFLICT OF INTEREST

Dr Bierut and the spouse of Dr Saccone are listed as inventors on U.S. Patent 8,080,371, 'Markers for Addiction' covering the use of certain SNPs in determining the diagnosis, prognosis and treatment of addiction. Authors listed with the affiliation deCODE Genetics/AMGEN are employees of deCODE genetics/AMGEN. Although unrelated to this research, Dr Kranzler has been a consultant or advisory board member for Lundbeck and Indivior and is a member of the American Society of

Clinical Psychopharmacology's Alcohol Clinical Trials Initiative, which was supported in the last 3 years by AbbVie, Alkermes, Ethypharm, Indivior, Lilly, Lundbeck, Otsuka, Pfizer, Arbor and Amygdala Neurosciences. Dr Kaprio has consulted for Pfizer in 2012–2014 on nicotine dependence. The remaining authors declare no conflict of interest.

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Supplementary Information accompanies the paper on the *Molecular Psychiatry* website (<http://www.nature.com/mp>)